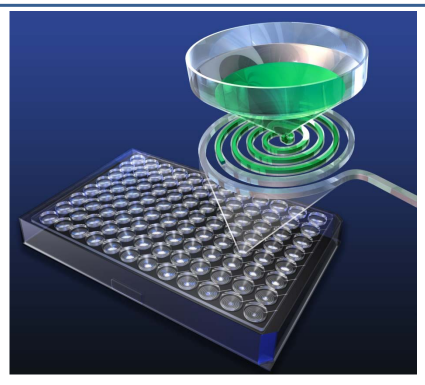
# **USER MANUAL SUPPLEMENT:**

Assay Solution for R&D Systems Mouse IL-23 DuoSet (R&D System Catalog DY1887)





Mouse IL-23 ELISA (R&D Systems Catalog# DY1887) has been successfully transferred to Optimiser™ microplate platforms to achieve the following key performance benefits.

• Sample Volume: 5 μl

• Assay time: Total assay time ~ 2 hours (~4.5 hour savings)

Assay reagents: 90% saving on antibody use
 150 assays for the cost of 15

Sensitivity/Range: Equivalent Sensitivity (39 – 2500 pg/ml)

• Potential to increase sensitivity to ~ 1.95 pg/ml

• Potential to achieve 3-log dynamic range: 39 – 28,400 pg/ml

## **Intended Use:**

This User Manual Supplement is intended to be used in conjunction with R&D Systems' Technical Data Sheet for the Mouse IL-23 DuoSet (Catalog Number DY1887) employed in this Optimiser™-based ELISA procedure. Please refer to the vendor's instructions for material storage, preparation, and concentration information.

The procedure described in this User Manual Supplement is intended as a starting reference for the investigator using the vendor's assay reagents with the Optimiser™ microplate system. Siloam has optimized this procedure using OptiBlock™ as blocking solution, as reagent diluent, and as the media to reconstitute the protein standard. Siloam has not evaluated this procedure for its applicability for analysis of tissue culture supernatants, serum, plasma, or other sample types. Use of this assay for the analysis of tissue culture supernatant, serum, or plasma samples may require further optimization of some assay parameters by the investigator (for example, standard curve and sample diluents) to achieve desired results.

It is expected that investigators following this assay procedure are familiar with the Optimiser™ microplate system. If you have not used the Optimiser™ microplate before, please order the Evaluation kit (Catalog# OPV-IL6) which provides a comprehensive overviewto the Optimiser™ microplate system. The Evaluation Kit guides the user through correct pipetting procedures for Optimiser™ microplates and contains all necessary materials and instructions for completing an illustrative human IL-6 assay. Investigators are strongly urged to familiarize themselves with the Optimiser™ microplate system before completing the procedure described in this document.

Please contact Siloam's tech support (<u>techsupport@siloambio.com</u>) for any questions regarding this procedure.

FOR RESEARCH USE ONLY
Not for Use in Diagnostic Procedures

# **MATERIALS:**

**Assay Reagents:** 

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Material <sup>1</sup>	Function	Vendor	Catalogue	Storage
			Number	
Anti-Mouse IL-23	Capture Ab			
Anti-Mouse IL-23 Biotin	Detection Ab			
r-Mouse IL-23	Standard	R&D Systems	DY1887	Per TDS
Streptavidin-HRP <sup>2</sup>	Streptavidin-			
	HRP			

# Optimiser™ Materials:

Material	Product Number	Storage
Optimiser™ plate (with holder) <sup>3</sup>	OPH-10	Room temp
OptiMax <sup>™</sup> buffer reagent pack (with substrate)	OMR-10-F	2 – 8 °C
Streptavidin-HRP for Optimiser™	OMR-HRP	2 – 8 °C

Other Material & Equipment:

Materials	Equipment
Polypropylene centrifuge tubes, 1.5 – 2 mL,	Fluorescence plate reader
snap-cap	
Pipet tips	Vortex mixer
KimWipes	Single channel pipettor(s)
Test tube rack	Multichannel pipettor
Reagent reservoirs (v-bottom)	

<sup>1</sup> 

<sup>&</sup>lt;sup>1</sup> Refer to the respective Technical Data Sheets (TDS) for storage, concentration and other relevant information.

 $<sup>^{2}</sup>$  DO NOT use SAv-HRP from assay reagent vendor. Use of OMR-HRP is strongly recommended.  $^{2}$ 

<sup>&</sup>lt;sup>3</sup> Optimiser™ plates and corresponding OptiMax™ buffer reagents are also available in 2-plate and 50-plate configuration.

#### **REAGENT PREPARATION:**

- Working concentrations of all materials should be prepared before beginning the procedure.
- 2. Capture antibody working solution:
  - a. Refer to the vendor's TDSfor the protein concentration of the stock capture antibody solution.
  - b. Prepare the capture antibody working solution by diluting the stock capture antibody to 8  $\mu g/mL$  in OptiBind<sup>M-F</sup>.

# 3. Lyophilized Standard:

- a. Reconstitute the lyophilized protein standard with OptiBlock™. Refer to the vendor's TDS for further directions and the concentration of the reconstituted standard.
- b. Prepare single-use aliquots by dispensingthe reconstituted standard to polypropylene tubes. Store at ≤ -70 °C.

#### 4. Standard Curve:

- a. Prepare standard 1 by diluting the reconstituted standard to 2500 pg/mL in OptiBlock™ (OB).
- b. Dispense 200 µL of standard 1 to well A1 of the polypropylene 96-well v-bottom plate.
- c. Dispense 100 μL of OptiBlock™ to each of the 7 wells immediately below well A1.
- d. Prepare serial two-fold dilutions of the standard by successive 100  $\mu$ L transfers through well G1.Change pipet tips after each transfer and mix the well contents 8-10 times by gently aspirating and dispensing the well contents. Do not transfer standard to well H1. Well H1 will serve as the assay blank (0 pg/mL).

	1	-				1
Α	200 μL Std 1	•			Α	Std 1
В	100 μL OB	•	190		В	Std 2
С	100 μL OB		IGO		С	Std 3
D	100 μL OB		199	/	D	Std 4
E	100 μL OB	•			E	Std 5
F	100 μL OB	•	180		F	Std 6
G	100 μL OB	•			G	Std 7
Н	100 μL OB	-	00		Н	Blank

#### Detection antibody:

- a. Refer to the vendor's TDS for the protein concentration of the stock detection antibody solution.
- b. Prepare the detection antibody working solution by diluting the stock detection antibody material to 0.2 µg/mL in OptiBlock™.
- 6. Horseradish-Peroxidase-Labeled Streptavidin (SAv-HRP):
  - a. Refer to the vendor's TDS or vial label for the dilution factor required to prepare the SAV-HRP working solution.
  - b. Prepare the SAv-HRP working solution by diluting the stock material appropriately in OptiBlock™.

#### 7. OptiGlow™ working solution:

- a. Prepare the OptiGlow™ substrate working solution by combining OptiGlow™- A, OptiGlow™- B, and OptiGlow™- C in proportions of 50:50:1 parts, respectively.
- Note: Prepare the working substrate solution no more than 30 minutes before reading the plate.

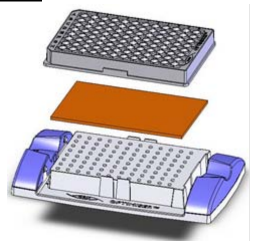
## Use of OptiBlock™:

This method was developed using Siloam Biosciences' OptiBlock™ as blocking agent following the coating of the microfluidic reaction chamber with capture antibody and as diluent for the standards, detection antibody, and SAv-HRP. The intent in developing this method was to demonstrate the ease of transitioning R&D Systems' Mouse IL-23 DuoSet from their intended use in conventional ELISAs to an Optimiser™-based ELISA format. Use of this method for the analysis of tissue culture supernatant, serum, plasma, or other sample types may require further optimization of some assay parameters by the investigator to achieve desired results; for example, standard curve and sample diluents.

Siloam Biosciences has developed OptiMax™ Standard Diluent for use in the analysis of cell culture supernatants. This product has been incorporated in Siloam Biosciences' commercially available OptiMax™ ELISA Kits.

#### DISPENSING MATERIALS TO THE OPTIMISER™ PLATE:

- Optimiser™ assembly: Assemble the Optimiser™ plate, pad, and holder as illustrated.
  - a. Position the holder on the lab bench with the Optimiser™ logo facing the user.
  - b. The absorbent pad must be positioned with the plasticcovered surface facing the holder.
  - Position the pad and plate on the holder surface and push down firmly until the plate snaps into position.



- 2. Reverse pipetting: Introduction of bubbles to the Optimiser™ wells will compromise method performance by occluding the microchannel. To avoid introducing bubbles, always use the reverse pipetting technique when delivering materials to an Optimiser™ well.
  - a. Beginning with the pipettor's operating button in the ready position, depress the operating button to the <u>second stop</u> (See figure in step c below).
  - b. Immerse the pipet tip in the liquid to be transferred. Aspirate the liquid by releasing the operating button returning it to the <u>ready position</u>.
  - c. Dispense the liquid to the Optimiser™ well by depressing the operating button to the <u>first</u> stop. Ensure that the pipet tip is touching the well surface as the liquid is dispensed.

	Pipetting Step			
Ready Position	1	2	3	4
First Stop		<b>†</b>	1	<b>†</b>
Second Stop	<b>\</b>			

- 3. <u>Transferring reagents, standards, and samples to the Optimiser™ plate:</u>Due to the short incubation times, it is critical that antibodies, standards, samples, SAv-HRP, and substrate are transferred from their source to the Optimiser™ wells quickly (< 1 minute) but accurately. To accomplish this:</p>
  - a. First, dispense the materials to a polypropylene 96-well v-bottom plate.
  - b. Then, using a multichannel pipettor, transfer the materials from the polypropylene vbottom plate to the Optimiser™ wells as illustrated in the figure on the next page.

#### Optimiser Washes:

Optimiser™-based ELISAs use a unique "flush" step rather than the traditional, and laborious, "wash" step used in conventional ELISAs. To flush, the user simply dispenses OptiWash™ into the Optimiser™ well. The wash buffer "flushes" the used reagent/sample from the microchannel into an absorbent pad beneath the plate. The Optimiser™ "flush" is equally effective as the traditional washes.

	1	2	3	4	5	6	7	8	9	10	11	12
Α	Std 1	Samp	Samp									
		1	9									
В	Std 2	Samp 2	Samp 10									
С	Std 3	Samp	Samp									
C	Jta 3	3	11									
D	Std 4	Samp	Samp									
	0.15	4	12									
E	Std 5	Samp 5	Samp 13									
F	Std 6	Samp	Samp									
'		6	14									
G	Std 7	Samp	Samp									
		7	15									
Н	Blank	Samp 8	Samp 16									
	← 5 µL of standard, sample, and blank are transferred from individual wells of polypropylene v-bottom plate to duplicate cells of Optimiser™.											
	1	2	3	4	5	6	7	8	9	10	11	12
Α	Std	1;	Sample #:	1	Sample i	#9						
В	Std	2;	Sample #2	2	Sample #10							
С	Std	3;	Sample #3 Sample #11									
D	Std	4;	Sample #4	4	Sample #12		Shade	d cells i	not used	d in this	assay	
E	Std	5;	Sample #!	5	Sample #13							
F	Std	6;	Sample #	6	Sample #14							
G	Std	7;	Sample #	7	Sample #15							
Н	Blar		Sample #8	3	Sample i	¥16						
	(0 pg/	mL)										

↑ Optimiser<sup>™</sup> plate to which standards, samples, and blank will be dispensed

PROCI	EDURE:
2.	() Dispense 5 µL of the working capture antibody solution to the appropriate
	number of wells in the Optimiser™ plate. Incubate 10 minutes at room
	temperature (RT).
3.	() Following the incubation, dispense 5 μL OptiWash™ to each well. Incubate
	10 minutes at RT.
4.	() Following the wash step, dispense 5 μL OptiBlock™ to each well. Incubate
	10 minutes at RT.
5.	() Following the block step, dispense 5 µL standard or sample to each well.
	Incubate <u>20</u> minutes at RT.
6.	() Following the sample incubation, dispense 5 μL OptiWash™ to each well.
	Incubate 10 minutes at RT.
7.	() Following the wash step, dispense 5 μL of the detection antibody working
	solution to each well. Incubate 10 minutes at RT.
8.	, , , , , , , , , , , , , , , , , , , ,
	well. Incubate 10 minutes at RT.
9.	( <u></u>
	each well. Incubate 10 minutes at RT.
10	0. () Following the SAv-HRP incubation, dispense 30 μL OptiWash™ to each well.
	Incubate 10 minutes at RT.
13	L. () Immediately following step 10, dispense 30 μL OptiWash™ to each well (for
	a second 30 μL wash). Incubate 10 minutes at RT.
12	2. ()Dispense 10 μL OptiGlow™ working solution to each well. Incubate 15
	minutes at RT.
	a. () Observe the plate periodically during this incubation. When the
	substrate solution has drained from all wells, first remove the plate from
	the holder and then remove the pad from the plate.
	b. () Wipe the bottom of the plate thoroughly with a KimWipe™ or
1.	similar laboratory tissue.  3. ( ) Read the plate using an excitation wavelength of 528/20 nm and an
13	emission wavelength of 590/35 nm.
	CHIISSION WAVEICHKIN OF JOZO JIIII.
a/	) This space is provided as a simple way of documenting the completion of each
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step.

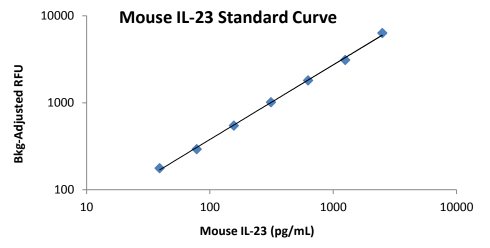
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## **Calculations:**

- 1. Calculate the mean background signal (RFU).
- 2. Subtract the mean background signal from the individual standard and sample values.
- 3. Calculate the mean background-adjusted signal for each standard and sample.
- 4. Prepare a standard curve by plotting the concentration of the standard on the x-axis and the background- adjusted signal on the y-axis using a 4-parameter curve fit.
- 5. Interpolate the sample concentrations from the standard curve. Calculate the final concentration after applying the sample dilution factor if applicable.

# Typical Data:

The standard curve illustrated below was generated using the method, reagents, and equipment specified in this procedure.



Additional technical assistance is available under the Technical Support tab on the Siloam Biosciences web site (<a href="http://www.siloambio.com/">http://www.siloambio.com/</a>).

- Material Safety Data Sheets (MSDS)
- Using Optimiser™ Immunoassay Microplate Video
- Optimiser™ User's Guide
- Reader Settings
- Quick Reference Guide
- Frequently Asked Questions
- Application Notes

Two additional videos appear under the Technology tab of the web site.

- Optimiser™ Principles of Operation
- Running an Assay with Optimiser™

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QuantaRed™ substrate is supplied by Thermo Fisher Scientific Inc.



# **Better Immunoassays Through Innovative Microfluidics**

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